



## Ultrashort Time-to-Echo Magnetic Resonance Imaging at 3 T for the Detection of Spondylolysis in Cadaveric Spines: Comparison With CT

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**Abstract:** **OBJECTIVES** The objective of this study was to compare the diagnostic performance and confidence of conventional, optimized, and ultrashort time to echo (UTE) magnetic resonance (MR) protocols for detection of simulated lumbar spondylolysis in human cadavers. In addition, we sought to demonstrate the feasibility of the UTE technique in subjects with and without spondylolysis. **MATERIALS AND METHODS** Four human lumbar spine specimens with 46 individual pars interarticularis were randomly left intact ( $n = 26$ ) or received experimental osteotomy ( $n = 20$ ) using a microsurgical saw to simulate spondylolysis. The specimens were imaged using a computed tomography (CT) scan along with 3 "Tiers" of MR protocols at 3 T: Tier 1, conventional lumbar MR protocol; Tier 2, optimized conventional protocol consisting of a sagittal oblique spoiled gradient recall echo and axial oblique T1 and short tau inversion recovery sequences; and Tier 3, a sagittal UTE MR sequence. Two blinded readers evaluated the images using a 4-point scale (1 = spondylolysis certainly absent, 2 = probably absent, 3 = probably present, 4 = certainly present) at each individual pars. For each imaging protocol, diagnostic performance (sensitivity, specificity, and area under the receiver operating characteristic curve, using the surgical osteotomy as the reference) and confidence were assessed and compared using the McNemar test. Furthermore, 2 human subjects were imaged with the conventional and UTE MR protocols to demonstrate feasibility in vivo. **RESULTS** Diagnostic performance was moderate for Tiers 1 and 2, with a moderate sensitivity (0.70 to 0.75) and high (1.00) specificity. In contrast, CT and Tier 3 UTE MR imaging had both high sensitivity (1.00) and specificity (1.00). The sensitivities of CT or Tier 3 were statistically greater than Tier 1 sensitivity ( $P = 0.041$ ) and neared statistical significance when compared with Tier 2 sensitivity ( $P = 0.074$ ). Area under the receiver operating characteristic curve was also significantly greater for CT and Tier 3 (each area = 1.00), compared with the areas for Tier 1 (0.89,  $P = 0.037$ ) or Tier 2 (0.873,  $P = 0.024$ ). Diagnostic confidences of CT or Tier 3 were much greater than other Tiers: Both Tiers 1 and 2 had a large percentage of uncertain ( $>60\%$ ,  $P < 0.001$ ) or wrong interpretations ( $>10\%$ ,  $P < 0.001$ ), unlike CT or Tier 3 (0% uncertain or wrong interpretations). Preliminary in vivo UTE images clearly depicted intact and fractured pars. **CONCLUSIONS** Our study demonstrated that the detection of pars fractures using a single sagittal UTE MR sequence is superior in performance and confidence to conventional and optimized MR protocols at 3 T, whereas matching those from CT evaluation. Furthermore, we demonstrated the feasibility of in vivo application of the UTE sequence in subjects with and without spondylolysis.

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results in very low contrast between the cortical bone and the pars defect, and an overall hypointense appearance masking the defect. Novel MR techniques, collectively referred to as ultrashort time to echo (UTE) sequences, have overcome this issue using TE of 1 millisecond or less<sup>19,20</sup> and acquiring sufficient MR signal from the scar tissue to detect spondylolysis.

We hypothesize that UTE technique is capable of detecting pars fractures as accurately as CT. The objective of this study was to compare the diagnostic performance and confidence of conventional, optimized, and UTE MR protocols for detection of simulated lumbar spondylolysis in human cadavers. In addition, we sought to demonstrate the feasibility of the UTE technique in subjects with and without spondylolysis.

MATERIALS AND METHODS

The cadaveric study part was exempted from institutional review board (IRB) approval. The in vivo study part was IRB-approved. All procedures performed in this study involving human participants were compliant with the regulations of the Health Insurance Portability and Accountability Act. The human participants gave written informed consent for study participation, collection of data, and use of prototype pulse sequences.

Study Population

Cadaveric Specimens

Four randomly selected human cadaveric lumbar spines obtained as a block of tissue provided by the U. C. San Diego Anatomical Material Program was used for our study. Twenty-three vertebral segments (vertebra T12–L5 in most cases) equaling 46 individual pars interarticularis from 4 different cadaveric spines (n = 4 donors; 3 women, 1 man; mean age, 54 ± 18 years) were available. The spine specimens included the paraspinal muscles and soft tissue structures, and did not have a history of back surgery.

Human Subjects

One 45-year-old male subject (height, 196 cm; weight, 102 kg) and one 46-year-old male subject (height, 170 cm; weight, 61 kg) with chronic lower back pain were retrospectively included in the ancillary in vivo part of our study. These patients were part of a broader imaging study on lower back pain, referred by a sports medicine physician (S. A.). No reference standard (eg, CT) was available for the UTE sequence (ie, index test) of these patients.

Experimental Pars Fractures

From the 46 available pars interarticularis, 20 were selected randomly using a computer random number generator, to be fractured experimentally. Remaining 26 pars were left intact. Cadaveric spines were thawed and consecutively operated using a standard midline posterior approach by a board-certified neurosurgery fellow with expertise in spine surgery (S. F.) with 7 years of experience. During the procedure, each pars interarticularis was exposed by gentle surgical preparation creating a continuous soft tissue flap on each side to ensure adequate adaptation of the paraspinal soft tissue structures. Visual evaluation of each pars interarticularis ensured that no preexisting structural defect of the pars was present. Experimental fractures were created (n = 20 pars), using a surgical oscillating saw (Stryker System 6; Stryker Corp, Kalamazoo, MI) with a 0.9- to 1.0-mm thick microsurgical bone blade (SO6–115 and SO6–131) similar to the technique published by Kepler et al.<sup>21</sup> Our experimental pars fractures would fall between stage II “early spondylolysis” characterized by a hairline fracture and stage III “progressive spondylolysis” characterized by a wide pars defect, according to the spondylolysis classification by Ralston et al.<sup>22</sup> Photo documentation was done after the fractures were performed. To ensure filling of the experimental fracture gaps with a material featuring similar MR signal characteristics as scar tissue, we tested 30 different materials including agarose (A9539; Sigma-Aldrich, Carlsbad, CA) in different concentrations. The candidate materials were filled in small vials and embedded in a slab of bovine muscle as a surrogate for paraspinal muscle. Of the candidates, 8% (by weight) agarose gel was selected as most suitable. Each fracture gap was filled with ~1 mL of 8% agarose solution applied with a syringe with an 18-gauge needle. After cooling and solidification of the agarose gel, the soft tissue structures were meticulously reattached using a deep and superficial suture technique. Subsequently, to prevent air accumulation in the adjacent soft tissue structures, each specimen was tightly wrapped in transparent film and vacuum-sealed. The surgical osteotomies served as the reference standard of the presence of pars fracture.

Magnetic Resonance Imaging

All MRI was performed on a 3 T GE MR Discovery 750 scanner (GE Healthcare, Milwaukee, WI) with a standard spine array coil on the following day after the experimental fractures were surgically induced. Three “Tiers” of MR protocols for the lumbar spine were created: (1) conventional, (2) optimized conventional, and (3) novel UTE MR sequences.

Tier 1, conventional MR protocol constituted of a standard lumbar MR protocol (Table 1). Tier 2, optimized conventional MR protocol

TABLE 1. MR Sequence Parameters of Tiers 1–3

	Sequence	TR, ms	TE, ms	Reconstruction Matrix	Slice, mm	FOV, cm	BW (±)	FA, degree	Voxel Dimension	Scan Time, min
									(xyz, mm)	
Tier 1	Sagittal SE T1	2297	24	256 × 256	3	24	62.5	90	0.94 × 0.94 × 3	2:34
	Sagittal SE T2	4618	102	256 × 256	3	24	62.5	90	0.94 × 0.94 × 3	2:14
	Axial SE T1	774	13	256 × 256	3.5	16	62.5	90	0.63 × 0.63 × 3.5	2:59
	Axial SE T2	9733	102	256 × 256	3.5	16	62.5	90	0.63 × 0.63 × 3.5	2:07
	Sagittal STIR	6466	60	256 × 256	3	24	31.25	90	0.94 × 0.94 × 3	3:07
Tier 2	Axial oblique SE T1	2297	24	256 × 256	3	24	62.5	90	0.94 × 0.94 × 3	2:34
	Axial oblique STIR	14938	60	256 × 256	3.5	16	31.25	90	0.63 × 0.63 × 3.5	3:38
	Sagittal SPGR	27	7	256 × 256	2	24	20.8	30	0.94 × 0.94 × 2	6:43
Tier 3	Sagittal 3D Cones	44.3	0.05	256 × 256	2	24	125	2	0.94 × 0.94 × 2	3:12

For the axial oblique sequences of Tier 2, the axial imaging plane was sloping down toward rear (Fig. 2E, dashed line H) since the pars is oriented obliquely to all 3 orthogonal planes. An oblique sagittal multiplanar reconstruction for the right and the left side was performed for the SPGR sequence.

SE indicates spin echo; 3D Cones, name of a particular UTE sequence; TR, time to repetition; TE, time to echo; FOV, field of view; BW, bandwidth; FA, flip angle; xy, in-plane; z, through-plane.



constituted of conventional MR sequences using imaging planes that are optimally aligned to the plane of the pars interarticularis according to previously published studies.<sup>11,15</sup>

Tier 3, UTE MR protocol constituted of a single sagittal 3D Cones sequence. All protocol specifications are summarized in Table 1, no additional planning or reconstruction time was required for the sagittal 3D Cones sequence. The 3D Cones sequence uses a unique k-space sampling trajectory that samples data along cone surfaces in 3D.<sup>23</sup> No postprocessing of the UTE dataset was used for the readout. Axial UTE images were also obtained for illustration purposes in some cases, but they were not used for the readout. The used UTE sequences are prototype pulse sequences.

For the ancillary in vivo part of the study, lumbar MRI of human subjects comprised the same sequences used in Tiers 1 and 3. No major changes to the protocol were needed, as the cadaveric study utilized MR sequences and coils that were readily translatable.

## CT Imaging

After the experimental fractures were surgically induced, the spine specimen were scanned using a 256-MDCT scanner (Revolution CT; GE Healthcare, Milwaukee) with 120 kV and 100 mA and a slice thickness of 0.625 mm with a reconstruction diameter of 50 cm and a matrix of 512 × 512. The fracture gaps were measured on sagittal reformatted images perpendicular to the lower side of the fracture line using RadiAnt DICOM Viewer software (Medixant, Poznan, Poland) with a slice thickness of 0.625 mm by a radiologist (C. B. C.) who did not participate in the readout.

## Detection of Spondylolysis in Images of Cadaveric Specimens

The assessment of spondylolysis detection of the vertebral segments between T12 and L5 was performed independently by 2 blinded, board-certified radiologists with fellowship training in musculoskeletal radiology. Reader 1 (R1) (P. S.) and reader 2 (R2) (T. F.) had 7 and 5 years of experience, respectively. The RadiAnt DICOM Viewer software (Medixant, Poznan, Poland) was used for assessment and provided the use of cross-reference lines and image leveling.

To reduce recall bias, we first waited 4 weeks after MRI before evaluating Tier 1 images, waited 2 weeks before reading Tier 2 images, and waited another 2 weeks before reading Tier 3 images, randomizing the order of samples between Tiers as well as readers. For this study, spondylolysis was detected as a continuous discontinuity in the cortical shell and cancellous bone in individual pars. For each Tier of images, the diagnostic confidence for the presence of spondylolysis was assessed using a 4-point scale: 1 (spondylolysis certainly absent), 2

(spondylolysis probably absent), 3 (spondylolysis probably present), 4 (spondylolysis certainly present), a scoring system used in previous studies by Lee et al.<sup>24</sup> and Jaeger et al.<sup>25</sup> For Tiers 1 and 2 where multiple sequences were evaluated, each reader noted which particular sequence was the most informative and decisive, if the reader found the pars to be positive for a fracture.

## Statistics

Interreader agreement for the detection of spondylolysis of each individual pars interarticularis was calculated using linearly weighted Cohen kappa ( $\kappa$ ) analysis where coefficient of 0.21 to 0.40 indicated fair, 0.41 to 0.60 moderate, 0.61 to 0.80 good, and greater than 0.81 indicated an excellent agreement.<sup>26</sup> For subsequent analyses, in case of a disagreement, a consensus reading was performed and eventually the score from the senior reader was used.

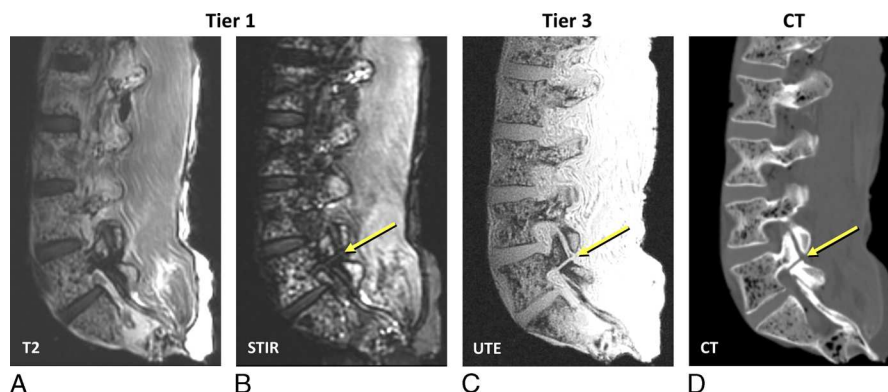
For each Tier, the diagnostic performance (sensitivity, specificity, and receiver operating characteristic area under the curve) and confidence (uncertain or wrong diagnosis) for the detection of spondylolysis was calculated using the surgical osteotomies as the reference standard. The sensitivity and specificity values were compared between Tiers, using McNemar statistic separately for sensitivity in samples positive for pars defect, and for specificity in samples negative for pars defect.<sup>27</sup> Area under the curve values were also compared<sup>28</sup> between Tiers. Diagnostic confidence values were also compared between Tiers using McNemar statistic. For statistical analysis, commercially available software was used (Systat 10; Systat Software, San Jose, CA). Statistical significance was set at a 2-tailed *P* value below 0.05.

## RESULTS

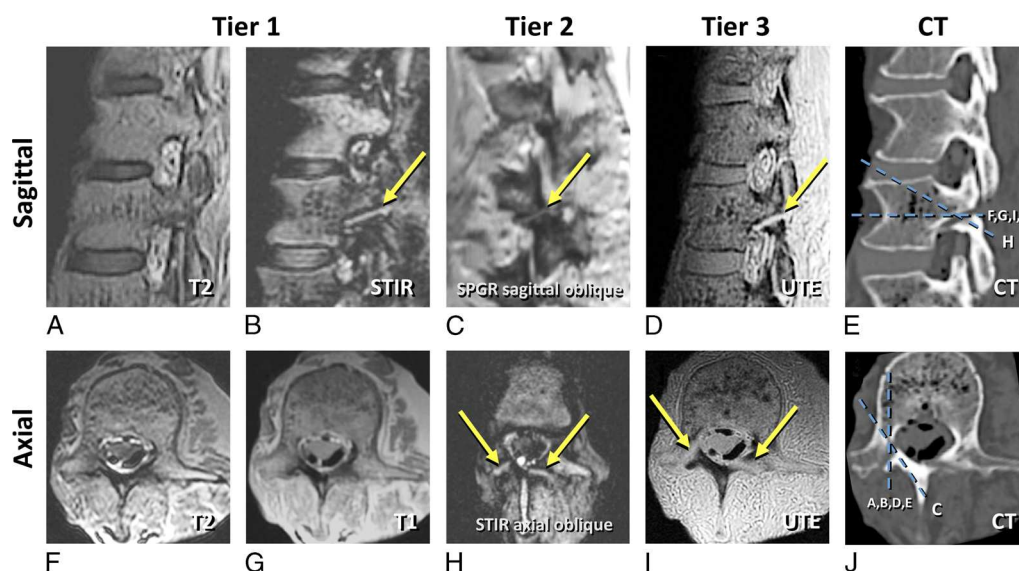
### Cadaveric Study

On the basis of the randomization, 20/46 (43%) pars interarticularis were selected for experimental fracture and 26/46 (57%) pars were selected to be left intact. The mean width of the experimental fracture gaps was  $1.2 \pm 0.3$  mm (range, 1.0–1.9 mm), only slightly wider than the thickness of the saw blade. Approximate reading time per specimen for each Tier was Tier 1 = 4 minutes, Tier 2 = 3 minutes, AND Tier 3  $\leq$  1 minutes.

Using conventional sequences of Tier 1, the pars fractures were usually not visible on axial and sagittal T1 and T2 (Fig. 1A) images, whereas the sagittal short tau inversion recovery (STIR) images often revealed the gap as a linear medium signal intensity (Fig. 1B, arrow). In general, evaluation of pars fractures on axial images (Fig. 2, F and G) was challenging due to varying oblique angles of the fractures to all 3



**FIGURE 1.** Right-sided imaging of cadaveric spine from a 42-year-old female donor, showing typical sagittal images of an experimental pars fracture at L5. On the conventional T2 images (A), the experimental pars defect on the right L5 level is not visible. The STIR sequence (B) shows the pars defect to some extent (arrow). In contrast, both the UTE (C) and the CT (D) images clearly depict the very thin experimental pars fracture, ~1 mm in width.



**FIGURE 2.** Imaging of cadaveric spine from a 56-year-old woman, showing representative images from all Tiers. Tier 1 (A, B, F, G), Tier 2 (C, H), Tier 3 (D, I), and CT (E, J) images are shown. Among the Tier 1 images, pars defect created on L1 level was not visible on sagittal T1 (not shown), sagittal T2 (A), axial T2 (F), and axial T1 (G) images, but it was moderately visible on sagittal STIR image (B, arrow). The rather horizontal orientation of the pars fracture, in combination with a slice thickness of 3.5 mm for the axial T2 (F) and T1 (G) sequences, hampered the detection of pars defects considerably. Among the Tier 2 images, sagittal oblique SPGR revealed the fracture at the neck of the “Scotty Dog” sign (C, arrow), and axial oblique STIR revealed the bilateral fractures with reduced partial volume artifact (H, arrows). Axial oblique T1 images (not shown) were less effective at revealing the fractures due to low signal intensity at the gap. In comparison, sagittal UTE (D) and CT (E) images clearly depicted the pars fracture (arrow) as a linear discontinuity. The pars fractures are also clearly visible in axial UTE (I, arrows) and CT (J) images but the margins of the defect are soft due to the partial volume artifact. These axial UTE and CT images were acquired for illustration purposes only and not used in the readout. The reference lines for the respective imaging planes are shown in the CT images (E, J).

orthogonal planes, as well as a complex anatomy involving close proximity of pars to facet joints (Fig. 2E).<sup>29</sup>

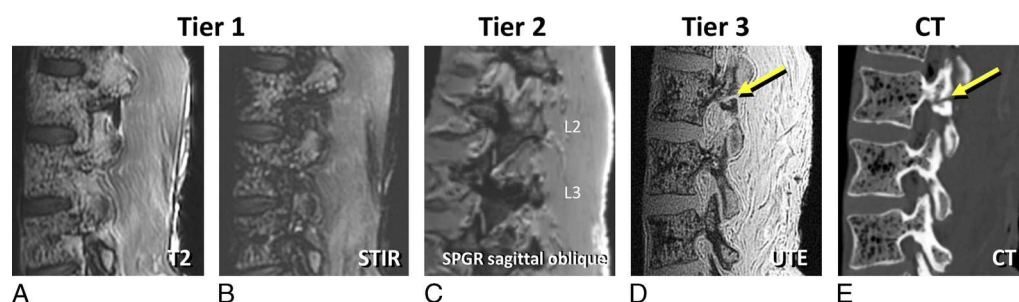
In Tier 2, by tilting the sagittal imaging plane or reformatting (Fig. 2J, dashed line C), sagittal oblique spoiled gradient recall echo (SPGR) images (Fig. 2C) revealed classic “Scotty Dog” sign that featured pars fracture at the neck of the dog (Fig. 2C, arrow). Similarly, by tilting the axial imaging plane (Fig. 2E, dashed line H) more orthogonal to the fracture, Tier 2 axial oblique STIR images (Fig. 2H) also made it easier to detect the pars fractures. In Tiers 1 and 2, the sagittal STIR and the sagittal oblique SPGR, respectively, were the most decisive sequences.

Although the optimized sequences of Tier 2 were an improvement over conventional sagittal and axial sequences of Tier 1, Tier 3 UTE images allowed the most facile detection of pars fractures. All pars defects were as clearly and confidently visible on sagittal UTE images (Figs. 1C, 2D, arrows) as they were on CT images (Figs. 1D, 2D). The similarities between UTE and CT in image characteristics (albeit

inverted) and the visibility of pars fractures due to the increased contrast between bone and fracture gap were striking. In a few difficult cases where the pars fractures were not visible on the conventional (Fig. 3, A and B) and optimized conventional (Fig. 3C) MR protocols, the single sagittal UTE sequence revealed the fractures with ease (Fig. 3D, arrow).

Interreader agreement was good for conventional Tier 1 ( $\kappa = 0.777$ ; confidence interval [CI], 0.647–0.907) and optimized Tier 2 ( $\kappa = 0.847$ ; CI, 0.734–0.960), and excellent for CT or Tier 3 ( $\kappa = 1.00$ ; CI, 1–1), suggesting an easier and straight-forward read for CT and Tier 3 UTE images compared with other Tiers.

Diagnostic performance (Table 2) was moderate for Tiers 1 and 2, with a moderate sensitivity (0.70 to 0.75) and high (1.00) specificity. In contrast, CT and Tier 3 UTE MRI had both high sensitivity (1.00) and specificity (1.00). The Tier 3 sensitivity was statistically greater than Tier 1 sensitivity ( $P = 0.041$ ), and nearly significantly greater than Tier 2 sensitivity ( $P = 0.074$ ). There was no difference in specificity



**FIGURE 3.** Imaging of cadaveric spine from a 42-year-old woman, showing a difficult case where the pars fractures were not detected on Tier 1 and 2 images. On the conventional sagittal T2 (A) and STIR (B) images (Tier 1), as well as on the sagittal oblique SPGR (C) images (Tier 2), the right-sided pars fracture at the L2 level was not detected during the readout. In contrast, on the sagittal UTE (D) and CT (E) images, a thin fracture (arrows) was easily detected with high diagnostic confidence.

TABLE 2. Diagnostic Performance

Pars Fracture	Tier 1: Conventional				Tier 2: Optimized				CT or Tier 3: UTE			
	Present		Absent		Present		Absent		Present		Absent	
	Certainly	Probably	Probably	Certainly	Certainly	Probably	Probably	Certainly	Certainly	Probably	Probably	Certainly
Present (n = 20)	5	9	6	0	3	12	3	2	20	0	0	0
Absent (n = 26)	0	0	19	7	0	0	16	10	0	0	0	26
Sensitivity			0.70*				0.75†				1.00	
Specificity			1.00				1.00				1.00	
AUC ROC			0.89*				0.87*				1.00	

Summary of the diagnostic performance of each MR Tier and CT images for pars fracture detection using the surgical osteotomies as the reference standard. The sensitivity was moderate (0.70 to 0.75) for Tiers 1 and 2. In contrast, CT and Tier 3 MR had higher sensitivity values (1.00). The Tier 3 sensitivity was statistically higher than Tier 1 sensitivity ( $P < 0.05^*$ ), and nearly significantly higher than Tier 2 sensitivity ( $P = 0.074^\dagger$ ). There was no difference in the specificity values between the imaging modalities. The area under the receiver operating characteristic curve (AUC ROC) was statistically greater for CT and Tier 3 compared with Tiers 1 and 2 (each  $P < 0.05^*$ ).

between the Tiers. Area under the receiver operating characteristic curve (Table 2) was also significantly greater for CT and Tier 3 (each area = 1.00), compared with the areas for Tier 1 (0.89,  $P = 0.037$ ) or Tier 2 (0.873,  $P = 0.024$ ).

More importantly, the diagnostic confidence of Tier 3 was much greater than the other Tiers. Table 3 shows the percentage of uncertain but correct (ie, spondylolysis probably absent or probably present, when a correct diagnosis is made) and wrong reads for each Tier. It can be seen that both Tiers 1 and 2 each have statistically higher ( $P < 0.001^{**}$ ) percentages of uncertain (>60%) and wrong (>10%) diagnoses, compared with CT or Tier 3.

In Vivo Study

In 2 subjects with low back pain, Tier 1 sagittal conventional T2 (Fig. 4, A and D) and Tier 3 UTE (Fig. 4, B and E) MR images were obtained. The axial UTE (Fig. 4, C and F) images were acquired for illustration purposes only and not used in the readout. In one subject, intact lumbar pars were well-depicted on the sagittal and axial UTE images (Fig. 4, B and C). In the second subject, suffering from a progressive nonradicular low back pain, the UTE sequences accurately revealed spondylolysis (Fig. 4, E and F, arrows) on L5 level of the male patient, which was not visible on the conventional T2 image (Fig. 4D) due to scar tissue within and near the pars defect. This subject had a wide, stage IV spondylolysis, according to Ralston et al<sup>22</sup> with an early anterolisthesis.

DISCUSSION

Early diagnosis of spondylolysis is crucial in young individuals to increase the likelihood of nonsurgical bone healing and to avoid non-union with potential spondylolisthesis in bilateral cases.<sup>30,31</sup> In adults featuring almost exclusively long-standing terminal spondylolysis, diagnosis of spondylolysis is important to ascertain the cause of back pain and to aid planning of spinal fusion surgery as appropriate.

The UTE sequence used in our study was able to match diagnostic performance of the CT, whereas outperforming conventional and optimized conventional protocols significantly. On standard MR sequences, short T2 tissues such as cortical bone and osteofibrous scar tissue surrounding the pars defect unequivocally remain dark,<sup>32</sup> whereas on UTE sequences, these structures provide contrast. It should be noted that UTE technique is able to acquire MR signal from cortical bone,<sup>33</sup> the MR signal from other soft tissues (including scar tissue) is markedly higher, leading to an appearance of dark bone surrounded by high signal intensity. These basic differences in MR signal characteristics provide the foundation for an improved detection of spondylolysis using UTE sequences. The sagittal UTE sequence is also practical in clinical routine requiring a scanning time of only 3 minutes and could therefore be added to a conventional MR protocol easily.

The sensitivity and specificity of our conventional and optimized lumbar MRI protocols for spondylolysis detection were similar to those reported in previous studies. A recent meta-analysis published by Dhouib et al<sup>12</sup> from 2017 included 1122 pars of young adults and children and showed a combined sensitivity of 81% (54 to 95% CI) and a combined specificity of 99% (98 to 100% CI), which is comparable to the results of our conventional (Tier 1) lumbar MR protocol. In studies that have investigated the diagnostic performance of a protocol similar to our optimized conventional (Tier 2) MR protocol, they showed a sensitivity of 86% to 92% and a specificity of 92% to 100%.<sup>10,34</sup> The slightly lower sensitivity of the optimized conventional MR protocol in our study could be explained by susceptibility artifacts introduced by small amounts of air and metal wear debris possibly introduced during sample preparation. Such artifacts were visible on a few SPGR images. Fortunately, UTE sequences are less prone to susceptibility artifacts of this kind.<sup>35,36</sup>

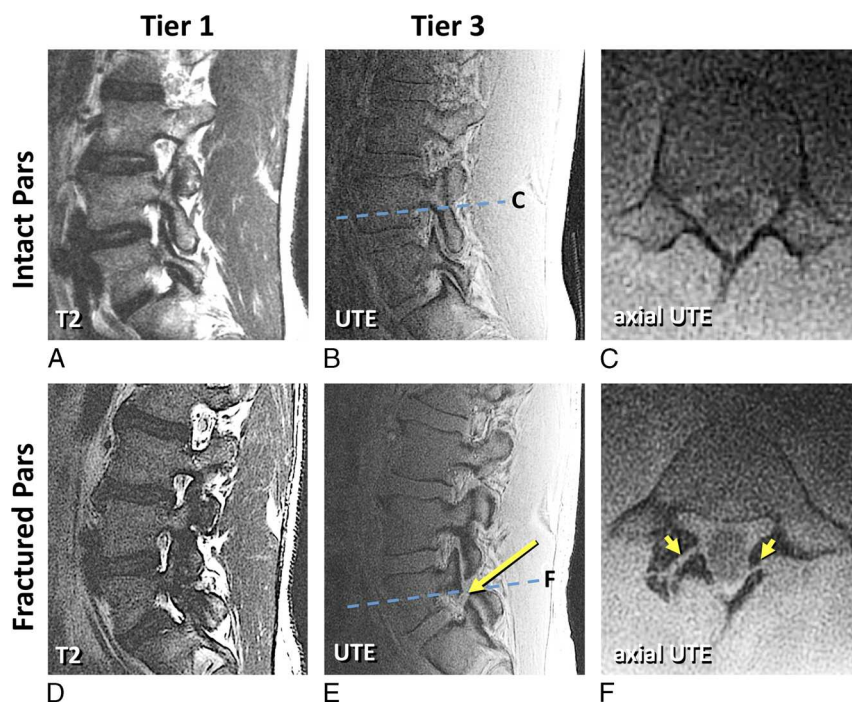
We have also demonstrated the feasibility of translating the UTE technique in vivo in a preliminary manner. In our human subjects with and without spondylolysis, image quality was similar to the specimen study, which was designed with clinical translation in mind. In future

TABLE 3. Diagnostic Confidence

Pars Fracture	Tier 1		Tier 2		CT or Tier 3	
	Uncertain	Wrong	Uncertain	Wrong	Uncertain	Wrong
All cases (n = 46)	61%**	13%**	61%**	11%**	0%	0%

Diagnostic confidence of imaging modalities expressed as the percentage of uncertain but correct reads (ie, spondylolysis probably absent or probably present, when a correct diagnosis is made) and wrong diagnoses. Results are divided according to the surgical osteotomy reference standard into “present” (pars fracture is present), “absent” (pars fracture is absent), and all cases. Notably, Tiers 1 and 2 have statistically higher ( $P < 0.001^{**}$ ) percentages of uncertain (>60%) and wrong (>10%) diagnoses, compared with CT or Tier 3.





**FIGURE 4.** In vivo imaging of subjects without (A–C) and with (D–F) spondylolysis. In a 46-year-old male subject (A–C), the right-sided L4 pars was intact on the sagittal T2 (A) as well as on the sagittal UTE (B) and axial UTE (C) image. In images of a 45-year-old male subject with bilateral terminal spondylolysis at L5 (D–F), sagittal T2 image (D) could not depict the fracture. In comparison, the corresponding sagittal (E) and axial (F) UTE images clearly depicted the pars defects (arrows), indicating the feasibility of in vivo application of the UTE sequence in subjects with and without spondylolysis.

studies, the capacity of UTE technique to diagnose spondylolysis in adults and ultimately in the pediatric population should be further investigated. Although the availability of our particular UTE sequence may not be widespread, currently other sequences capable of ultrashort echo time can run on clinical MR systems from almost every major manufacturer (eg, encoding time reduction with radial acquisition [PETRA] and zero echo time sequences) and have been increasingly used in medical imaging studies.<sup>37,38</sup> For example, the recently published study by Breighner et al<sup>39</sup> found great merit of zero echo time sequences for bone imaging of the shoulder.

It should be clarified that our study does not address all stages of spondylolysis, including very early stages. According to Ralston et al,<sup>22</sup> there are 4 successive stages of spondylolysis. Our experimental fractures would fall between stage II “early spondylolysis” characterized by a hairline fracture and stage III “progressive spondylolysis” characterized by a wide pars defect. The other spondylolysis stages, prespondylolytic stage I “pars stress reaction” and stage IV “terminal spondylolysis” characterized by nonunion and sclerosis, could not be created experimentally, because these are dependent on edematous bone marrow response and bone remodeling in a living tissue. However, edematous bone marrow changes can be accurately detected using conventional fat-suppressed fluid-sensitive sequences,<sup>3,15,16,22,40</sup> and it is likely that the diagnostic performance of all Tiers will be increased for even larger fractures of terminal stage spondylolysis (such as that seen in our subject in Fig. 4, E and F, arrows). Although our experimental fractures with ~1 mm width were fairly thin, it remains to be determined if even thinner hairline fractures could also be detected with UTE techniques.

Our study has further limitations. First, due to the small number of available specimens, we created an artificially large number of fractures (20 of 46 pars, at 2 or more levels in all specimens). This frequency exceeds the prevalence of approximately 5% in the adult population and approximately 50% in adolescent athletes with persistent back pain,<sup>3</sup> usually occurring only at a single level at L5.<sup>1</sup> Second, our surgical preparation inherently constituted an artificial condition.

However, we meticulously applied a technique previously published by Kepler et al,<sup>21</sup> using even thinner oscillating saw blades of only 0.9 to 1 mm, whereas preserving the paravertebral soft tissue structures as good as possible. Nonetheless, the CT proved that the morphology and orientation of the experimental fractures were similar to natural fractures.<sup>41</sup> Third, we did not include other types of MR sequences that may better visualize bone detail such as 3D Turbo spin echo MRI, susceptibility-weighted MRI,<sup>42</sup> or “black bone” gradient echo MRI.<sup>43,44</sup> It would be useful to evaluate and compare the performance of these sequence for spondylolysis image. Fourth, to prevent air and fluid accumulation, the delicate fractures were filled with agarose gel. As described, we systematically identified the 8% agarose gel as the best candidate featuring MR signal characteristics similar to scar tissue that is frequently located within and in proximity to pars defects. Lastly, our study was performed at 3 T only and therefore the results may or may not be transferable to lower field strength scanners. There are confounding issues such as susceptibility artifacts from cortical bone at 3 T,<sup>45</sup> which may vary with the field strength.

In conclusion, our study demonstrated that the detection of pars fractures using a single sagittal UTE MR sequence is superior in performance and confidence to conventional and optimized MR protocols at 3 T, whereas matching those from CT evaluation. Furthermore, we demonstrated the feasibility of in vivo application of the UTE sequence in subjects with and without spondylolysis. We believe that UTE sequences offer a great potential for radiation-free bone imaging and will play an increasing role in patient care in the future.

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